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(54) Title: A DISEASE RESISTANCE GENE FROM MAIZE AND ITS USE FOR DISEASE RESISTANCE, AS A SELECTABLE MARKER AND AS A GENE IDENTIFICATION PROBE

(57) Abstract

The HMI gene in maize confers race-specific resistance to the pathogen, Cochliobolus carbonum. We have used transposon mutagenesis to tag, clone, and characterize several HMI alleles. The gene can be used as a selectable marker in conjunction with the toxin produced by C. carbonum.

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A DISEASE RESISTANCE GENE FROM MAIZE AND ITS USE FOR DISEASE RESISTANCE, AS A SELECTABLE MARKER AND AS A GENE IDENTIFICATION PROBE

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TECHNICAL FIELD

This is a continuation-in-part of prior copending application No. 07/995,658, filed December 15, 1992, now abandoned. This invention relates to the isolation of a gene which controls resistance to both a fungus and a fungal disease toxin and its use to confer resistance to fungal disease, as a selectable marker and as a probe to identify homologous disease resistance genes in maize and other crops.

15 BACKGROUND OF THE INVENTION

Disease resistance genes are defined as Mendelian factors that cosegregate with the resistance trait. The gene <u>HM1</u>, which controls resistance to <u>Cochliobolus carbonum</u> Nelson race 1 was among the first disease resistance genes to be described. The disease caused by <u>C. carbonum</u> race 1 can be devastating, resulting in yield losses of 80% or more due to plant death and grain mold. The dominant allele, <u>Hm1</u>, and the duplicate factor, <u>Hm2</u> are the only disease resistance genes that are known to be fixed at a high frequency in maize germplasm.

Since the discovery of a race-specific compatibility factor that is produced by the fungus, the disease caused by <u>C. carbonum</u> race 1 has been the subject of detailed study. This compatibility factor permits the fungus to infect certain genotypes of maize that would otherwise be resistant. The role of <u>HM1</u> in providing resistance to the fungus is a function of its ability to cause reduced sensitivity to the compatibility factor. Further studies established that the presence of the compatibility factor confers the same race-specificity (for hm1/hm1 corn) on the oat pathogen, <u>Cochliobolus victoriae</u>. The structure of this compatibility factor, HC-toxin, is known but the mode of action remains to be elucidated. Recently, an

enzyme that inactivates HC-toxin has been identified in extracts from maize. The enzyme, HC-toxin reductase (HCTR), is detectable only in extracts from resistant (Hm1) genotypes. This establishes HCTR activity as the biochemical phenotype of Hm1.

DISCLOSURE OF THE INVENTION

It has now been determined that the Hml gene can be used in conjunction with HC-toxin in a selectable marker system for use in maize transformation. When the cloned 10 gene is linked to appropriate regulatory sequences for expression in plant cells and cotransformed into maize cells along with another quantitative or qualitative trait which is not selectable, it confers upon transformants a resistance to HC-toxin by virtue of the production of The cells can continue to grow on medium containing the isolated HC-toxin. Nontransformed cells do not express the HCTR and are rapidly killed by the toxin the pathogen produces. The net effect is a tissue culture containing only transformed cells which can then be regenerated by known methods to form transformed shoots and even whole plants.

It has also been discovered that this gene can be inserted using conventional transformation methods into the genomes of maize varieties which lack the gene and are susceptible to disease caused by <u>C carbonum</u>. Resulting transformants are resistant to the disease.

It has further been discovered that the HM1 resistance gene is one of a family of homologous disease resistance genes in maize and other crops. Accordingly, the present invention also encompasses methods for identification of homologous disease resistance genes by using this gene or a fragment thereof as a probe.

Materials and Methods

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Genetic Materials and Propagation

Mutator stocks used in this work were obtained from Dr. D. Robertson, Iowa State University. Ac/Ds stocks used were obtained from D. I. Greenblatt, University of Connecticut. The methodology of HM1 mutant isolation was

performed in the manner previously published by one of us [S.P. Briggs, Curr. Top. Plant Biochem. Physiol., 6:59 (1987)].

Inbred <u>HM1</u> allele designations are: Pr, <u>hm1-1</u>; K61, <u>hm1-2</u>; P8, <u>Hm1-A</u>; Pr1, <u>Hm1-Pr1</u>; B79, <u>Hm1-B79</u>; 4Co63, Hm1-4C063; Pioneer PHV12, Hm1-PHV12.

Experimental Approach

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Transposon-tagging is known to be a reliable tool for isolating genes from maize. The basic tenet of tranposon tagging is that DNA rearrangements (i.e., insertions or excisions) that are concomitant with genetic (phenotypic) changes define a causal relationship which identifies the rearranged DNA as the changed gene. Several independent events of this nature are accepted as proof of identity. As described herein, we have observed that 5 mutant alleles of HM1 are associated with insertions within a transcribed region (hm1-656::Mu1), hm1-1369::Mu3, 1062::dHbr, hm1-2355, hm1-1040::Spm), and 1 deletion allele Def(HM1)-1790 is associated with loss of the transcribed region. Finally, only the wild-type allele (Hm1-Pr1) produced a 1.3 kb mRNA.

transposable element family Mutator The is particularly effective for generating forward mutations. Germinal reversion of mutants back to wild-type is rare with Mutator-induced alleles. Somatic reversion has been observed, but the small somatic sectors typical of Mutator cannot be identified for traits that are not cell autonomous or are complex, such as disease development. Therefore, to obtain multiple genetic events that were coincident with rearrangements of HM1, strategy based upon independent forward mutations rather than reversions was used.

To identify co-segregation between <u>HM1</u> and a restriction fragment, the segregating progeny were first classified by examining their DNA on Southern blots. The blots were hybridized with probes for RFLP loci that flank the locus; PIO200644 and PIO200044 (obtained from D.

Grant, Pioneer Hi-Bred International, Inc.) map 5 cM proximal and distal to HM1, respectively (2; unpublished observations). Only progeny that inherited alleles at both loci from the same parent were used in the analysis. This approach permitted the identification of progeny which inherited the 10 cM block of chromosome 1 bounded by By grouping together progeny that loci. inherited either the mutant allele, or the recessive tester allele with which the mutant allele was paired, and classes with each then comparing the two restriction fragments that were common to one progeny class and absent from the other were identified. method established linkage between restriction fragments and the 10 cM block that contains HM1.

To directly determine the linkage between HM1 and a restriction fragment, linked fragments were cloned and used to prepare DNA hybridization probes. Probes were then hybridized to a Southern blot of DNA from 60 progeny of the backcross K61/Prl X K61. The backcross progeny were scored for inheritance of the HM1 alleles by inoculating them with conidia of the C. carbonum race 1 strain SB111, and for inheritance of the alternate alleles detected by the DNA probes. A comparison of the patterns of inheritance revealed the linkage relationships between the probes and HM1.

The RFLP loci described above, plus an RFLP locus that is detected by the probe NPI429, were used to identify the progenitor of the mutant alleles. We obtained NPI429 from T. Helentjaris, Native Plants, Inc.

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Isolation of DNA and blot hybridization

Total DNA from the leaf tissue of maize, <u>Sorghum</u> (Pioneer inbred P285), and <u>Coix</u> (<u>Lachrymae jobi</u>) and seedling tissue of <u>Arabidopsis thaliana</u> L. (<u>Landsberg erecta</u>) was isolated by the urea extraction method as described in S. L. Dellaporta, J. Wood, J.B. Hicks, <u>Plant Mol Biol Rep</u>, 1, 18 (1983). Southern blots were prepared as described by Athma et al, <u>Genetics</u>, 128, 163 (1991).

For RFLP analysis, DNA was transferred to nylon membranes (MSI from Fisher) and the hybridizations were performed as described above but without the addition of formamide. Probes were made from gel-purified DNA fragments and labeled by random priming (Amersham). The Mul specific probes was an internal 650 bp Aval/BstEII fragment isolated from pA/B5; the plasmid was provided by L. Taylor, Stanford University. The Mul specific probe was the internal HindIII/XbaI fragment of a clone that was obtained from V. Chandler, University of Oregon. The Spm probes were pBx1 and pXS2.3 obtained from K. Cone, Brookhaven National Laboratory.

Isolation of RNA and Northern Blotting

Total RNA from 5 to 6 day-old etiolated seedlings was isolated by the guanidine thiocyanate method [P. Chomczynski and N. Sacchi, Anal Biochem 162, 156 (1987)]. Poly(A) + RNA was enriched using the polyATtract mRNA isolation system of Promega. Samples (~15 ug) of poly(A) + RNA were denatured using formaldehyde, fractionated in a 1.3% agarose gel, and blotted onto Hybond-N by standard techniques (). DNA probes were radiolabeled by random priming and the blots were hybridized and washed as described.

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Genomic Cloning

DNA isolated from homozygous mutant seedlings was digested with SstI or XhoI and the appropriate DNA fragments (as judged from the Southern blots) were purified by preparative gel electrophoresis electroelution into dialysis tubing. This purified DNA was ligated to preannealed SstI or XhoI cut arms from the bacteriophage vector 1 sep6-lac5 obtained from Martienssen, Cold Spring Harbor Laboratory. Packaging into Gigapack Gold (Stratagene) and screening of the libraries were carried out according to the manufacturers instructions. All clones were subcloned into Bluescript

(Stratagene) and maintained in the SURE strain (Stratagene) of E. coli.

PCR (Polymerase Chain Reaction) and Sequence Analysis

For analysis of DNA from the resistant progeny of the heterozygote, hm1-656::Mul/hm1-1369::Mu3 homologous to sequences on each side of the insertion sites were used for PCR amplication. The primer sequences were: 5' CTG CTC ATG ACT CAT ATC AGG CGG TAG C 3' AND 5' 10 GAC CAG CCG ACG CAG CAG CCC CGC CTT C 3'. PCR conditions were as described by Perkin Elmer-Cetus, except that the reactions were performed in 20% glycerol. Reactions were heated to 94 C for 3 min, then cycled 40 times for 1 min at 94 C, 2 min at 65 C, and 2 min at 72 C; and finally extended for 15 min at 72C. The gel-purified PCR products were reamplified and directly sequenced using synthetic sequencing primers.

HM1 Mutants Derived from Mutator Stocks

Several mutant alleles of HM1 were recovered from Mutator element stocks, 4 of which were selected for Criteria for selection included DNA further study. hybridization data that ruled out pollen contamination as the source of the mutant alleles and a low frequency of susceptible progeny appearing within a family. were used to distinguish between the mutant alleles and the standard recessive alleles with which they had been paired. Segregating progeny were classified inheritance of mutant alleles by use of the RFLP probes.

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hm1-656::Mul

Progeny produced by fertilizing plants genotype y wx gl1 Hm1-B79 Mutator (designated 81-82-9537 Mu² per se by D. Robertson) with pollen from the hybrid, 35 K61/Pr, resulted in the recovery of 2 susceptible plants out of 253 progeny. Both susceptible plants were found to be carrying the mutant allele, hm1-656::Mu1, indicating that they arose as the result of a small somatic sector on

the ear. Pollen from both plants was used to fertilize ears on the Pioneer inbred, PHV12.

The *656 probe was hybridized to a Southern blot of DNA from 4 different homozygous mutants. Classification using RFLP loci indicated that each of the mutant alleles was derived from the B79 inbred parent (Hm1-B79). observation of polymorphisms showed that DNA rearrangement took place in the cloned region concomitant with the generation of the mutations. With the enzyme used, hm1-1062::dHbr produces a fragment identical to B79. be detected using other enzymes.

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hm1-1369::Mu3

The hm1-1369::Mu3 allele was recovered as a single susceptible plant out of 230 progeny. The progeny were produced by pollinating plants of the genotype y wx gll Hm1-B-79 Mutator (designated 81-82-9537 Mu2 per se by Dr. Robertson) with the hybrid, K61/Pr (hm1-2/hm1-1).

The heterozygote, hml-1369::Mu3/hml-2, was self-pollinated and the progeny were classified using RFLP loci. Hybridization of a Mu3 probe revealed a co-segregating band. A 4.6 kb SstI fragment co-segregated with the hml-1369::Mu3 allele. The signal intensity difference caused by 2 doses of the allele, when homozygous, permitted hml-1369::Mu3 to be scored as a

codominant marker, which increased the mapping resolution. No recombinants were observed in a sample of 60 progeny. The 4.6 kb fragment, which contained Mu3 plus flanking DNA, was excised from the gel and cloned into the SstI site of 1 sep6-lac5 subcloned into Bluescript SK+, restriction mapped, and used to prepare a probe (designated *1369) from the DNA that flanked the Mu3 insertion.

The *1369 probe detected the same polymorphisms that had been revealed by the *656 probe, indicating that the Mu1 and Mu3 elements had inserted into the restriction fragment. This conclusion was confirmed by comparison of the restriction maps of the 2 clones. anomaly was revealed with SstI in which the hm1-656::Mul DNA gave a 3.2 kb fragment when hybridized with the *656 probe but a 1.0 kb fragment when the *1369 probe was used; other enzymes failed to detect this anomaly. DNA sequence analysis showed that the Mul element had created an SstI site upon insertion into HM1.

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An effort was made to recover resistant progeny from the Mu alleles. The heterozygote, hm1-656::Mu1/hm1-1369::Mu3 was fertilized using pollen from the inbred Pr (hm1-1). From 500 progeny, 1 resistant plant was Examination of DNA from 10 susceptible recovered. siblings of the resistant plant confirmed that all three alleles were segregating as expected. Examination of DNA from the resistant plant revealed a progenitor-sized fragment from B79 (designated Hm1-B79R) plus the hm1-1 allele from Pr. This plant was self-DNA from its progeny was examined using pollinated. probes *1369, PIO200644, PIO200044, and NPI429. progeny were tested for susceptibility to infection by results confirmed that inoculating with SB111. The resistance was conferred by the Hm1-B79R Homozygous Hm1-B79R progeny were identified. The site of insertion of Mul/Mu3 was amplified by PCR and sequenced with no difference observed between the new allele and the B79 progenitor.

hm1-1062::dHbr

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Progeny produced by pollinating the hybrid, K61/Pr (hm1-2/hm1-1), with plants of the genotype y wx gl1 Hm1-B79 Mutator (designated 81-82-9539 Mu² per se by Dr. Robertson) resulted in the recovery of the hm1-1062::dHbr allele from 1 of 2 susceptible plants out of 483 progeny.

Hybridization with probes specific for Mul, Mul, Mul, Mul, and Mul failed to identify a fragment that cosegregated with hml-1062::dHbr. Hybridization with the *656 probe detected a polymorphism between hml-1062::dHbr and the progenitor allele, Hml-B79. We cloned a 3.1 kb XhoI fragment from the hml-1062::dHbr mutant into 1 sep6-lac5, using *656 as a probe. Restriction mapping confirmed the B79 structure with the exception of a small (approximately 400 bp) insertion into the SstI-XhoI fragment at the 3'-end of the clone. The identity of this insertion element has not been determined, but it lacks homology with Ac, Ds1, Ds2, and Spm/En internal sequence probes and with the Mu terminal inverted repeat.

Def(HM1-1790)

Progeny produced by pollinating plants of the genotype $y \times gl1 \times$

Def(HM1)-1790 allele The displays an aberrant transmission pattern. A Def(HM1)-1790/hm1-1 heterozygote was self-pollinated and the progeny were characterized using the flanking RFLP loci. Only 14 of 56 progeny were found to have inherited the Def(HM1)-1790 chromosome. When a heterozygote was fertilized using pollen from a wild-type stock (SB509), 8 of 25 progeny inherited the Def (HM1)-1790 chromosome. When the heterozygote was used as the pollen source to fertilize a wild-type inbred (W23), none of 32 progeny inherited the Def(HM1)-1790 The results show that the Def(HM1)-1790 chromosome.

allele (or chromosome) was not transmitted through the pollen and was only poorly transmitted through the egg. Such a pattern of transmission is typical of a chromosomal deficiency. The *1369 probe did not hybridize with DNA from the Def(HM1)-1790 allele, confirming the presence of a deletion and showing that the cloned region lies within the deletion. Test crosses with br2, which maps within 0.1 cM of HM1, produced only wild-type progeny. Likewise, both PIO200644 and PIO200044 detected 2 alleles in progeny that inherited Def(HM1)-1790. Therefore, the deletion cannot encompass more than 5 cM of the chromosome, being bounded by br2 and one of the RFLP loci.

An HM1 mutant derived from an Ac/Ds stock

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An allele, Hm1-1040::Spm, was recovered from the inbred 4Co63; the P-VV allele (Ac inserted into the P1 gene) had been backcrossed into this version of 4Co63. This allele was selected for study because it appeared to arise as a large tassel sector. The hybrid K61/Pr (hm1-2/hm1-1) was fertilized using pollen from the P-VV/P-(Hm1-4Co63/Hm1-4Co63), inbred 4Co63 designated 610417(X) by I. Greenblatt (personal communication). Seventeen males were used to produce 32 ears. Only 2 ears bore susceptible progeny and both were derived from male The cross 741-11 X 741-9 plant 16 plant number 16. yielded 47 susceptible progeny out of 323. The cross 741-13 X 741-9 plant 16 yielded 16 susceptible progeny out of The results are best explained by a single mutagenic event that occurred during tassel development, giving rise to many gametes bearing the same mutation.

Genetic tests showed that <u>Ac</u> had not transposed in the <u>hm1-1040::dSpm</u> mutant; <u>Ac</u>, <u>Ds1</u>, and <u>Ds2</u> probes failed to identify a restriction fragment that cosegregated with hm1-1040::dSpm. Examination of DNA from the hm1-1040::dSpm mutant by hybridization with the *1369 probe revealed an 11.0 kb SstI restriction fragment that was 6 kb larger than the Hm1-4Co63 progenitor allele. This fragment was cloned into 1 sep6-lac5. Hybridization of

various probes to DNA from the clone identified the insertion as an Spm/En homologous element.

Natural Variation

The Hm1 parental alleles from which the mutants described in this report were derived provide complete resistance to C. carbonum race 1 throughout development of the plant. In contrast, the Hml-A allele, found in the inbred P8, provides partial resistance that increases as the plant develops. Examination of DNA from P8 by hybridization with the ± 1369 probe showed that the cloned region is duplicated. The duplicate segregates independently of HM1. The relationship between the pattern of expression and the duplication has not been The ± 1369 probe hybridized well with DNA established. from the grasses Sorghum and Coix but poorly with DNA from Arabidopsis. Poly(A) + RNA from our mutants and from the inbreds K61 ($\underline{hm1-2}$) and Pr1 ($\underline{Hm1-Pr1}$) was blotted and hybridized with the ± 1062 probe. A 1.3 kb mRNA band was present only in the Prl lanes. The susceptible genotypes either had no detectable hybridizing mRNA or the size was aberrant. In all cases, the signal was extremely weak.

of the <u>HM1</u> gene has been cloned and sequenced, with genomic and cDNA sequences as shown in SEQUENCE I.D. No. 1 and SEQUENCE I.D. No. 2, respectively. Recessive alleles of the gene clearly contain homologous DNA (e.g., hm1-1 and hm1-2). This stands in contrast to the 2 fungal plant pathogen genes that have been cloned which control race specificity:: TOX2 confers race 1 type pathogenicity upon C. carbonum (Walton, personal communication) and avr9 confers race specific avirulence upon Cladosporium fulvum; both genes are missing in strains that lack their corresponding functions.

Resistance appears to differ from susceptible at the transcriptional level, at least among the small samples of alleles that were examined. This indicates that

susceptible genotypes do not possess an alternative form of <u>HM1</u> with specificity for a substrate other than HC-toxin. This result also suggests that the only function of <u>HM1</u> in young leaf tissue is to provide resistance, since the HM1 mutations are not pleiotropic.

Use in Disease Resistance

Plants

The preferred embodiment of this method involves inserting the HM1 gene into the genome of the plant in proper reading frame, together with transcription initiator and promoter sequences active in the plant. Transcription and translation of the gene under control of the regulatory sequences causes resistance to the disease caused by <u>C. carbonum</u>.

The plant must be a plant susceptible to infection 15 and damage by C. carbonum. These include corn (Zea mays). However, this is not to be construed as limiting, inasmuch as this species is among the most difficult commercial crops to reliably transform and regenerate, and this pathogen can also infect certain other crops. Thus, the 20 methods of this invention are readily applicable via conventional techniques to other plant species, if they are found to be susceptible to C. carbonum, including, without limitation, species from the genera Allium, Arabidopsis, Arachis, 25 Antirrhinum, Asparagus, Atropa, Avena, Beta, Brassica, Browallia, Capsicum, Cicer, Cicla, Citrullus, Citrus, Cucumis, Cucurbita, Datura, Daucus, Fragaria, Geranium, Digitalis, Fagopyrum, Glycine, Gossypium, Helianthus, Hordeum, Hemerocallis, Lactuca, . Lotus, Lycopersicon, Majorana, 30 Lens, Lolium, Manihot, Nicotiana, Medicago, Nasturtium, Oryza, Pelargonium, Persea, Petunia, Phaseolus, Pisum, Ranunculus, Raphanus, Ricinus, Saccharum, Secale, Senecio, Setaria, Solanum, Triticum, Spinacia, Trifolium, Bromus, Cichorium, Nemesia, Panicum, 35 Hyoscyamus, Linum, Onobrychis, Pennisetum, Salpiglossis, Sinapis, Trigonella, and Vigna.

Preferred plants that can be transformed according to the methods of this invention are cereal crops, including

maize, rye, barley, wheat, sorghum, oats, millet, rice, triticale, sunflower, alfalfa, rapeseed and soybean.

Numerous plant expression cassettes and vectors are well known in the art. By the term "expression cassette" is meant a complete set of control sequences including initiation, promoter and termination sequences which function in a plant cell when they flank a structural gene in the proper reading frame. Expression frequently and preferably contain an assortment restriction sites suitable for cleavage and insertion of any desired structural gene. In addition, the plant preferably cassette includes expression a constitutive promoter sequence at one end to cause the gene to be transcribed at a high frequency, and a poly-A recognition sequence at the other end for processing and transport of the messenger RNA. An example of such a preferred (empty) expression cassette into which the cDNA of the present invention can be inserted is the pPHI414 plasmid developed by Beach, et al. of Pioneer Hi-International, Inc., Johnston, Iowa, disclosed in the U.S. Patent Application No. 07/785,648, filed October 31, 1991, the disclosures of which are hereby incorporated herein by reference. Highly preferred plant expression cassettes will be designed to include one or more selectable marker genes, such as resistance or herbicide tolerance genes.

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By the term "vector" herein is meant a DNA sequence which is able to replicate and express a foreign gene in a host cell. Typically, the vector has one or more endonuclease recognition sites which may be cut in a predictable fashion by use of the appropriate enzyme. Such vectors are preferably constructed to additional structural gene sequences inparting antibiotic or herbicide resistance, which then serve as markers to identify and separate transformed cells. markers/selection agents include kanamycin, phosphonothricin, chlorosulfuron, hygromycin and methotrexate. A cell in which the foreign genetic

material in a vector is functionally expressed has been "transformed" by the vector and is referred to as a "transformant".

A particularly preferred vector is a plasmid, which is meant a circular double-stranded DNA molecule which is not a part of the chromosomes of the cell.

As mentioned above, genomic, cDNA or synthetic DNA encoding the HM1 gene may be used in this invention. vector of interest may also be constructed partially from a cDNA clone and partially from a genomic clone, etc. Genetic constructs are made which contain the necessary regulatory sequences to provide for efficient expression of the gene in the host cell. According to this invention, the genetic construct will contain (a) a first genetic sequence consisting of the HM1 gene, and (b) one or more regulatory sequences operably linked on either side of the structural gene of interest. Typically, the regulatory sequences will be selected from the group comprising of promoters and terminators. The regulatory sequences may be from autologous or heterologous sources.

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Promoters that may be used in the genetic sequence include nos, ocs, FMV and CaMV promoters.

An efficient plant promoter that may be used is an overproducing plant promoter. Overproducing promoters that may be used in this invention include the promoter of the small sub-unit (ss) of the ribulose-1,5biphosphate carboxylas from soybean (Berry-Lowe, et al., J. Molecular and App. Gen., 1:483-498 (1982)), and the promoter of the chlorophyll a-b binding protein. 30 two promoters are known to be light-induced, in eukaryotic plant cells (see, for example, Genetic Engineering of Plants, An Agricultural Perspective, A Cashmore, Pelham, New York, 1983, pp. 29-38, G. Coruzzi, et al., J. Biol. Chem., 258:1399 (1983), and P. Dunsmuir, et al., Molecular and App. Gen., 2:285 (1983)).

The expression cassette comprising the HM1 gene operably linked to the desired control sequences can be ligated into a suitable cloning vector. In general,

plasmid or viral (bacteriophage) vectors containing replication and control sequences derived from species compatible with the host cell are used. The cloning vector will typically carry a replication origin, as well as specific genes that are capable of providing phenotypic selection markers in transformed host cells. Typically, genes conferring resistance to antibiotics or selected However, in the context of this herbicides are used. invention, the HM1 gene can also serve as the selectable marker in the transformation process. After the genetic material is introduced into the target cells, successfully transformed cells and/or colonies of cells can be isolated by selection on the basis of survival in the presence of HC-toxin.

Typically, an intermediate host cell will be used in the practice of this invention to increase the copy number of the cloning vector. With an increased copy number, the vector containing the HM1 gene can be isolated in significant quantities for introduction into the desired plant cells. Host cells that can be used in the practice of this invention include prokaryotes, including bacterial hosts such as E. coli, S. typhimurium, and Serratia marcescens. Eukaryotic hosts such as yeast or filamentous fungi may also be used in this invention.

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The isolated cloning vector will then be introduced into the plant cell using any convenient technique, including electroporation (in protoplasts), retroviruses, microparticle bombardment, and microinjection into cells from monocotyledonous or dicotyledonous plants in cell or transformed plant tissue culture to provide containing as foreign DNA at least one copy of the plant expression cassette containing the HM1 gene. Using known techniques, protoplasts can be regenerated and cell or tissue culture can be regenerated to form whole fertile plants which carry and express the HM1 gene according to invention. Accordingly, a highly this embodiment of the present invention is a transformed maize plant, the cells of which contain as foreign DNA at least

one copy of an expression cassette of this invention containing the HM1 gene.

Finally, this invention provides methods of imparting resistance to diseases caused by <u>C. carbonum</u> to plants of a susceptible taxon, comprising the steps of:

- (a) culturing cells or tissues from at least one plant from the taxon,
- (b) introducing into the cells or tissue culture at least one copy of the expression cassette comprising the HM1 gene operably linked to plant regulatory sequences which cause the expression of the gene in the cells, and
- (c) regenerating disease-resistant whole plants from the cell or tissue culture. Once whole plants have been obtained, they can be sexually or clonally reproduced in such a manner that at least one copy of the sequence provided by the expression cassette is present in the cells of progeny of the reproduction.

By the term "taxon" herein is meant a unit of botanical classification of genus or lower. It thus includes genus, species, cultivars, varieties, variants, and other minor taxonomic groups which lack a consistent nomenclature.

It will also be appreciated by those of ordinary skill that the plant vectors provided herein can be incorporated into Agrobacterium tumefaciens, which can then be used to transfer the vector into susceptible plant cells, primarily from dicotyledonous species. Thus, this invention provides a method for imparting antimicrobial resistance in Agrobacterium disease and tumefaciens-susceptible dicotyledonous plants in which the expression cassette is introduced into the cells by infecting the cells with Agrobacterium tumefaciens, a plasmid of which has been modified to include a plant expression cassette of this invention.

35 Use as a Probe

This invention also provides methods of isolating disease resistance genes from maize and other species, by using the HM1 gene as a probe. It has been determined

that HM1 is homologous to a family of disease resistance genes in maize and other species such as Arabidopsis, and this gene provides a probe which can be used, hybridizing the HM1 gene with restriction digests of "target" genomic DNA, to count the disease resistance It can also be used, by hybridization genes therein. against a genomic library from the subject organism, to identify positive colonies which can then be cloned and sequenced. These techniques are well known and widely published, as seen by the published series Current Protocols in Molecular Biology, F. M. Ausubel, Ed., (Wylie 1987-89), which have disclosed such methods for the last eight years, and the disclosures of which are hereby incorporated herein by reference. The novel claimed herein is the use of the HM1 gene or a fragment 15 thereof in those methods as a highly selective probe for specific identification of disease resistance genes.

SEQUENCE LISTING

- (1) GENERAL INFORMATION:
 - (1) APPLICANT: Pioneer Hi-Bred International, Inc.
- (ii) TITLE OF INVENTION: A Disease Resistance Gene from Maize
 5 And Its Use For Disease Resistance, As A Selectable Marker And As A
 Gene Identification Probe
 - (iii) NUMBER OF SEQUENCES: 2
 - (iv) CORRESPONDENCE ADDRESS:
 - (A) ADDRESSEE: Pioneer Hi-Bred International, Inc.
- 10 (B) STREET: 700 Capital Square, 400 Locust

Street

- (C) CITY: Des Moines
- (D) STATE: Iowa
- (E) COUNTRY: United States
- 15 (F) ZIP: 50309
 - (v) COMPUTER READABLE FORM:
 - (A) MEDIUM TYPE: Diskette, 3.5 inch, 1.44 Mb storage
 - (B) COMPUTER: IBM Compatible
 - (C) OPERATING SYSTEM: MS-DOS
- 20 (D) SOFTWARE: Microsoft Windows Notepad
 - (vi) CURRENT APPLICATION DATA:
 - (A) APPLICATION NUMBER:
 - (B) FILING DATE:
 - (C) CLASSIFICATION:
- 25 (vii) PRIOR APPLICATION DATA:
 - (A) APPLICATION NUMBER:
 - (B) FILING DATE:
 - (viii) ATTORNEY/AGENT INFORMATION:
 - (A) NAME: Roth, Michael J., et al.
- 30 (B) REGISTRATION NUMBER: 29,342
 - (C) REFERENCE/DOCKET NUMBER: 0212R PCT
 - (ix) TELECOMMUNICATION INFORMATION:
 - (A) TELEPHONE: (515) 248-4895
 - (B) TELEFAX: (515) 248-4844
- 35 (2) INFORMATION FOR SEQ ID NO: 1:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 5198 base pairs
 - (B) TYPE: nucleotide

(C) STRANDEDNESS: double

(D) TOPOLOGY: linear (ii) MOLECULE TYPE: genomic DNA (A) DESCRIPTION: 5 (iii) HYPOTHETICAL: No (iv) ANTI-SENSE: NO (v) FRAGMENT TYPE : (vi) ORIGINAL SOURCE: (A) ORGANISM: Zea mays (B) STRAIN: 10 (C) INDIVIDUAL ISOLATE: (D) DEVELOPMENTAL STAGE: (E) HAPLOTYPE: (F) TISSUE TYPE: 15 (G) CELL TYPE: (H) CELL LINE: (I) ORGANELLE: (vii) IMMEDIATE SOURCE: (A) LIBRARY: (B) CLONE: 20 POSITION IN GENOME: (viii) (A) CHROMOSOME/SEGMENT: (B) MAP POSITION: (C) UNITS: (ix) FEATURE: 25 (A) NAME/KEY: (B) LOCATION: (C) IDENTIFICATION METHOD: (D) OTHER INFORMATION: (x) PUBLICATION INFORMATION: 30 (A) AUTHORS: (B) TITLE: (C) JOURNAL: **VOLUME:** (D) ISSUE: 35 (E) PAGES: (F) DATE: (G)

(H) DOCUMENT NUMBER:

- (I) FILING DATE:
- (J) PUBLICATION DATE:
- (K) RELEVANT RESIDUES IN SEQ ID NO:
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 1:
- GAATTCGTAT ATCAGTTTAC TGCATGTATA TTTTTTGCAT GCACATTGGA 50 TTTATGTTCG TAATTGCTAG ATTTTTACGT CTTCCATAAA ACTGTCCCTA 150 GTCTTAACTG CCAAATTTTT ATATCTGCCA TAAAATTGTC CCTAATTTCC 250 10 GTCGGCGTTC TTAGTTGGAC CGATATTAGG TTTTTTCAAT TATGCTACAC TGTCTAAATA TTTATGCGAC GTTGTATAAG AATTTGTGCT TTGTGTGACT CATGTCATCC AATTTTTTGT GCGCGTGCAG TGGAAAGGAC ATATAATCTA 400 AAATTTGCGT GCATGCAATG GAAACTCCTA CAATTTGCCT TAATTTTTAG 450 ATGTGCATAA AAATAGAAAT TGCATGCATA CGACTCCCAT ATTTTTCAGA 500 TCTGCCATAA AAATAGGATC GTAAATACAA CCCACCAAAC TATCAACCTC 550 TCATATTGGC TCGATATTTA TGTTTATAAC TGAGAGGTTT TATGCCTAAA 600 ACTTAAGGTA TTTACGCTCG AACCTGTAGA TGCATCCACC AGTGAACCAC 650 ATGCCAGATT TTTTACGTAG TATACCACAT TGCATATATA ACTACACCAT 700 GTTGCATAAT TTGTATAAGT ATATATCATA AACTGGTGCT AACGAGCCTA 750 20 GTTGCATGTC TGTTGAAGCG ATCATATATT TATACACGAA ATAAAGCATT 800 TAAATAATTT TTTTTTGTTT TCATGCATGC CAATCCATTT CCTTTCATCG 850 CACATCCTGC CATCCTAAAT TTGTGTGCAT GCATCAAGAA ACAGATTTTT 900 ACACAGTATA CTGAATTGAA TAAATACCTA CACCTTGTAA CATAATTAAT 950 ATCAGTATAT TATAAACTAG TCCTAACGAA TTTAGTTGTA TGGCTGTTGG 1000 TTCCATGCAT GCCAATCCAT TTCCTTTCAT CGCATATTCT TGTCATCTTA 1100 AATTTGTGCT CATGCAACAG GAAACAAATT TTTACGCAGT GTACCAAATT 1150 GCATAAACAC CTACACCGTG TTGCATAATT AGTATCGTTA TATAACATAA 1200 ACTAGTCTTG ACAAGCCTAT CTAAATGGGT TTTAGAGTGA TCTTATATTT 1250 30 ATGGAGGAAA TAAAGCATTA AATATGAATA TTTGTTTTCA TACATGACAA 1300 TTCATTTCCT TTTATCGTAC AAATTTTGCC ATCCTAATTT TGTGCACATG 1350 CAGCAGGTAA AAAACTTTTA CGTAGTGTAC CAAATTGCAT AAATATCTAC 1400 ACCGTGTAGC ATATTTAGTA TTAGTATATA TATCATAAAC TGGTCCTAAC 1450 GAGCCTAGCT GCATAGCTGT TGCAGCGATC CTATATTTAT GGAGAAAATG 1500 AATCATTTAA AAATAGAAAA AGTCAGAATC TTTTTTTTAA ÁACTTTACGC 1550 GTATGCAGGG ACACATGTTT GACTTATGCA TCCATTCTTT TTTTGCGAGC 1600 ATAAACGTGG GGTGGGGAGC GCACCCGACA AACTGGCTTG GGGAGTGAGT 1650 GGGCGCGCCG GCCTGGACCA GGTCGCAGGG GTGGTCGGCC TGGGCTTCCT 1700

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    CATATCCCTC GTCATGCCCA ACTGTCCTCC CGCCGTCGCA GAGAATTC 5198
     (2) INFORMATION FOR SEQ ID NO: 2:
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35

(1) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 1374 base pairs
- (B) TYPE: nucleotide
- (C) STRANDEDNESS: double
- (D) TOPOLOGY: linear

	(A)	DESCRIPTION:			
	(iii) HYPOTHETICAL: No				
	(iv) AN	TI-SENSE: NO			
5	(v) FRAG	MENT TYPE :			
	(vi) OR	IGINAL SOURCE:			
	(A)	ORGANISM: Zea mays			
	(B)	STRAIN:			
	(C)	INDIVIDUAL ISOLATE:			
10	(D)	DEVELOPMENTAL STAGE:			
	(E)	HAPLOTYPE:			
	(F)	TISSUE TYPE:			
	(G)	CELL TYPE:			
	(日)	CELL LINE:			
15	(I)	ORGANELLE:			
	(vii) IM	MEDIATE SOURCE:			
	(A)	LIBRARY:			
	(B)	CLONE:			
	(viii)	POSITION IN GENOME:			
20	(A)	CHROMOSOME/SEGMENT:			
	(B)	MAP POSITION:			
	(C)	UNITS:			
(ix) FEATURE:					
:	(A)	NAME/KEY:			
25	(B)	LOCATION:			
	(C)	•			
	(D)				
		LICATION INFORMATION:			
	(A)	AUTHORS:			
30	(B)	TITLE:			
		JOURNAL:			
	, ,	VOLUME: ISSUE:			
	(E)	PAGES:			
35	(E) (G)				
	(G) (H)				
	(I)				
•	(J)	FORTIGHTON DATE:			

(ii) MOLECULE TYPE: cDNA

(K) RELEVANT RESIDUES IN SEQ ID NO:

	(xi)	SEQUENCE DI	ESCRIPTION:	SEQ ID NO:	2:	
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5	GCTCCTGGCT	CGTCAGGAAG	CTCCTCGAGA	AAGGCTACAC	CGTCCACGCC	150
	ACCCTGCGGA	ACACCGGGGA	CGAGGCGAAG	GCGGGGCTGC	TGCGTCGGCT	200
	GGTCCCCGGC	GCGGCGGAGC	GGCTGCGGTT	GTTCCAGGCC	GACCTCTTCG	250
	ACGCCGCCAC	CTTCGCGCCG	GCGATCGCTG	GGTGCCAGTT	CGTCTTCCTC	300
	GTCGCCACGC	CATTCGGGCT	CGATAGCGCC	GGCTCCCAGT	ATAAGAGCAC	350
10	GGCGGAAGCT	GTGGTGGACG	CGGTGCGCGC	GATCCTCCGG	CAATGCGAGG	400
	AGTCCCGGAC	GGTGAAGCGA	GTGATCCACA	CAGCCTCCGT	AGCGGCCGCC	450
	TCGCCGTTGC	TGGAGGAGGA	GGTCTCCGCC	TCCGGCGTCG	GGTACAGAGA	500
	CTTCATCGAC	GAATCTTGTT	GGACTTCGCT	CAACGTTGAC	TATCCTCTCC	550
	GAAGCGCACA	CTTCGACAAG	TACATACTGT	CGAAGCTGCG	GTCAGAGCAG	600
15	GAGCTCCTGA	GCTACAACGG	CGGCGAGAGC	CCGGCGTTCG	AGGTGGTGAC	650
	CCTGCCGCTG	GGGCTCGTGG	CGGGCGACAC	GGTCCTCGGC	CGCGCCCCGG	700
	AGACGGTGGA	GAGCGCCGTG	GCGCCCGTGT	CCCGCAGCGA	GCCCTGCTTC	750
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	CGTGGACGAC	GTCTGCGACG	CGCTCGTCTT	CTGCATGGAG	CGGCGCCCCT	850
20	CCGTCGCCGG	CCGCTTCCTC	TGCGCCGCCG	CGTACCCGAC	GATCCACGAC	900
	GTGGTCGCCC	ACTACGCCAG	CAAGTTCCCT	CACCTCGACA	TCTTGAAAGA	950
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			AAGAAACGTG			
			TAGTCCACCT			
			CGATAATTTT	AGGCCTCTAA	AAAAAAAA	1350
30	AAAAAAAAA	AAAAAAAACT	CGAG			1374

WHAT IS CLAIMED IS:

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1. A DNA sequence coding substantially solely for the Hml gene of maize.

- 2. A DNA sequence having at least 90% translational homology to the sequence of SEQUENCE I.D. No. 1 or SEQUENCE I.D. No. 2.
- 3. An expression cassette containing the DNA sequence of claim 1 or 2 operably linked to plant regulatory sequences which cause the expression of the DNA sequence in plant cells.
- 4. A bacterial transformation vector comprising an expression cassette according to Claim 3, operably linked to bacterial expression regulatory sequences which cause replication of the expression cassette in bacterial cells.
- 5. Bacterial cells containing as a foreign plasmid at least one copy of a bacterial transformation vector according to Claim 4.
 - 6. Transformed plant cells containing at least one copy of the expression cassette of Claim 3.
- 7. A transformed plant comprising transformed cells according to claim 6.
 - 8. A method of identifying plant transformation using C. carbonum or the toxin produced by C. Carbonum as a phytotoxic marker, comprising the steps of:
- 25 a) culturing cells or tissues from a selected target plant,
 - b) introducing into the cell or tissue culture at least one copy of an expression cassette according to Claim 3,
 - c) introducing C. carbonum or the toxin it produces into the cell or tissue culture and
 - d) identifying transformed cells as the surviving cells in the cell or tissue culture.
- 9. A method of imparting resistance to diseases caused by C. carbonum to plants of a susceptible taxon, comprising the steps of:

a) culturing cells or tissues from at least one plant from the taxon,

- b) introducing into the cells or tissue culture at least one copy of an expression cassette comprising the
 5 HM1 gene operably linked to plant regulatory sequences which cause the expression of the gene in the cells, and
 - c) regenerating disease-resistant whole plants from the cell or tissue culture. Once whole plants have been obtained, they can be sexually or clonally reproduced in such manner that at least one copy of the sequence provided by the expression cassette is present in the cells of progeny of the reproduction.

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- 10. A method of identifying disease resistance genes which are homologous to the HM1 gene in the genome of a plant, comprising the step of identifying sequences which successfully hybridize with the HM1 gene or a fragment thereof.
- 11. A method according to Claim 10 wherein the sequences are identified by hybridizing a cDNA library or a restriction digest of genomic DNA from the plant with the HM1 gene or a fragment thereof.